

## Research Article

# Geochemical Fingerprinting of Rising Deep Endogenous Gases in an Active Hypogenic Karst System

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The hydrothermal caves linked to active faulting can potentially harbour subterranean atmospheres with a distinctive gaseous composition with deep endogenous gases, such as carbon dioxide (CO<sub>2</sub>) and methane (CH<sub>4</sub>). In this study, we provide insight into the sourcing, mixing, and biogeochemical processes involved in the dynamic of deep endogenous gas formation in an exceptionally dynamic hypogenic karst system (Vapour Cave, southern Spain) associated with active faulting. The cave environment is characterized by a prevailing combination of rising warm air with large CO<sub>2</sub> outgassing (>1%) and highly diluted CH<sub>4</sub> with an endogenous origin. The  $\delta^{13}\text{C}_{\text{CO}_2}$  data, which ranges from −4.5 to −7.5‰, point to a mantle-rooted CO<sub>2</sub> that is likely generated by the thermal decarbonation of underlying marine carbonates, combined with degassing from CO<sub>2</sub>-rich groundwater. A pooled analysis of  $\delta^{13}\text{C}_{\text{CO}_2}$  data from exterior, cave, and soil indicates that the upwelling of geogenic CO<sub>2</sub> has a clear influence on soil air, which further suggests a potential for the release of CO<sub>2</sub> along fractured carbonates. CH<sub>4</sub> molar fractions and their  $\delta\text{D}$  and  $\delta^{13}\text{C}$  values (ranging from −77 to −48‰ and from −52 to −30‰, respectively) suggest that the methane reaching Vapour Cave is the remnant of a larger source of CH<sub>4</sub>, which was likely generated by microbial reduction of carbonates. This CH<sub>4</sub> has been affected by a postgenetic microbial oxidation, such that the gas samples have changed in both molecular and isotopic composition after formation and during migration through the cave environment. Yet, in the deepest cave locations (i.e., 30 m below the surface), measured concentration values of deep endogenous CH<sub>4</sub> are higher than in atmospheric with lighter  $\delta^{13}\text{C}$  values with respect to those found in the local atmosphere, which indicates that Vapour Cave may occasionally act as a net source of CH<sub>4</sub> to the open atmosphere.

## 1. Introduction

Hypogene karstification is generally related to the rising of CO<sub>2</sub>- or H<sub>2</sub>S-rich fluids, and aggressiveness of the waters is obtained by cooling the fluids in the oxidation zone, not only in the water bodies but also in the air by condensation-corrosion processes. Caves form by the specific conditions of hypogene speleogenesis and that their active hydrogeochemical mechanisms (e.g., hydrothermal input, sulphuric acid, mixing corrosion, dissolution of evaporites, and dissolution in mixed sulphate-carbonate sequences [1–3])

can potentially harbour subterranean atmospheres with distinctive gas compositions. One example of this is when abiotic CO<sub>2</sub> and CH<sub>4</sub> gases are formed by chemical reactions that do not directly involve organic matter. This composition results from current activity or residual signs of degassing from gas-enriched groundwater or geothermal focus at depth. Consequently, this represents a mixture of multiple sources.

The migration of deep endogenous gases plays a key role in the formation of macroscopic void-conduit systems under hypogenic settings. Hypogenic karst regions are widely

distributed throughout the world [3]. However, there have been few studies addressing the gas composition of the subterranean atmospheres currently undergoing hypogene speleogenesis, and the few that exist have primarily focused on sulphidic caves (e.g., [4]). Air monitoring in the Frasassi Caves of Italy revealed a remarkable outgassing of both  $\text{CO}_2$  and  $\text{H}_2\text{S}$  from the groundwater, which varied seasonally from 1 to 8 ppm for  $\text{H}_2\text{S}$  and from 1500 to 5600 ppm for  $\text{CO}_2$ . In this system, the rapid gas exchange with the upper, non-sulphidic levels maintains a normal oxygen concentration [5]. Ascending warm water with  $\text{H}_2\text{S}$  and  $\text{CO}_2$  degassing, high salinity from dissolved chlorides, sulphides, high noble gas concentrations, and radioactive decay have been described as the main factors existing in hyperkarst phenomena along active deep-rooted faults [6]. Recently, it has been noted that the oxidation of  $\text{CH}_4$  influences the generation of sulphides during sulphuric acid speleogenesis in active systems [7]. This presents a new and valuable dataset of  $\text{CH}_4$  concentrations and stable-isotope ratios of hydrogen and carbon in  $\text{CH}_4$  found within the air of actively forming sulphidic caves.

Most hypogenic caves formed by the movement of thermal waters in phreatic settings are now located far above the water table and are thus no longer active. One exception to this pattern may be represented by caves associated with active faulting and geothermal areas where there are high concentrations and releases of upwelling fluids of endogenous origin, which enhance the processes and distinctive features of hypogene karstification [8–12]. As an example, venting of subcrustal  $\text{CO}_2$  has been described in a hydrothermal cave associated with an active, deep-rooted fault [13]. Recent works deal with the importance of seismic characterization of these systems since faults act as preferential migration routes of the ascending fluids that contribute to the formation of the hypogenic karstic system [14].

Vapour Cave (VC) in southern Spain represents a hypogenic system related to upwelling of deep endogenous fluids from an active faulting zone. The pathways and mechanisms that control the exchange of deep endogenous gases among atmosphere, soil, and subsurface reservoirs in VC have not been characterized to date. This study is aimed at characterizing the specific constraints imposed by an active hypogenic system such as that found in VC in the dynamics of deep-sourced gases (carbon dioxide and methane) with potential for release into the open atmosphere. Here, we provide insights into the behaviour of these deep endogenous gases in the upper vadose zone of karst terrains in active faulting zones. Isotope ratios of carbon and hydrogen imprinted in the rising geogenic gases ( $\text{CO}_2$  and  $\text{CH}_4$ ) in this hypogenic cave are used to investigate the sourcing and biogeochemical processes involved in the release, storage, and consumption of these gases into the upper vadose zone, and their interaction with the lower troposphere.

## 2. Geological Context and Hydrogeological Settings

VC is located within the Murcia Province of SE Spain (Figure 1), along the southern side of a small carbonates butte

in the village of Alhama de Murcia. This small butte is part of the *Sierra Espuña*, and it is directly related to the Alhama de Murcia Fault (AMF), a tectonically active, NE-SW-trending master fault with a left lateral strike-slip and a reverse component. The last earthquake attributed to this fault was in May of 2011, during which nine people were killed, and many rock-falls and ground cracks affected a total area of nearly 1000 km<sup>3</sup> around the epicentre zone [15]. Cave development affected Miocene (Lower Tortonian) conglomerates with rounded pebbles of carbonate and metamorphic rocks [16]. This unit is related to a postorogenic mantle of carbonate units structured by reverse faults, which have since changed to strike-slip tectonics. Convergence between Africa and the microplate of Iberia has uplifted the mountains of *Sierra de Carrascoy* and *Sierra Espuña* along a dominantly NE-SW trend.

The study zone of Alhama de Murcia is located within the Segura River basin (SE of Spain) (Figure 1). The hill where VC is located (namely, “Cerro del Castillo,” at 319 m a.s.l.) has been traditionally related with the quaternary alluvial deposits of the Low Guadalentin aquifer. However, the last studies [17] established a hydrological connection with the western carbonates belonging to the Santa Yechar aquifer (namely, Santa Yechar-Alhama aquifer), which are quite massive, of Triassic age, and with an average thickness of 150 m. Therefore, the hydric recharge primarily happens in southern foothills of *Sierra Espuña* mountains, where the main carbonate outcrops of the Santa Yechar aquifer are located (Figure 1). The superficial area of the Santa Yechar-Alhama aquifer is 58 km<sup>2</sup>.

Figure 2 summarizes the main geological units where this aquifer is embedded in the vicinity of VC. The main permeable unit of this aquifer is formed by 150 m thickness of Triassic black dolostones (Tr in Figure 2), from the Alpujarride Complex. On this unit and showing discordance appears 100 m of polygenic conglomerates of Tortonian age (G in Figure 2, host-rock of Vapour Cave) and a stratum of red sandstone and conglomerates of middle-lower Tortonian age (A in Figure 2). The impermeable units at the bottom correspond to Permo-Triassic phyllites (Pz in Figure 2), of the Intermediate Betic Units and Alpujarride Complex. At local scale, the aquifer is laterally sealed by marls of middle-lower Tortonian age, which form the relief of the northern mountains (*Sierra de la Muela*), and to the south with the late Miocene marls with gypsum located below the quaternary alluvial deposits (m and R in Figure 2, respectively).

At regional scale, the structure of this aquifer is determined by three mega nappes caused by an overthrust with south dip. This tectonic structure determines that the groundwater bodies of this aquifer are linked to the location of each one of these nappes, which are widely compartmentalised by an extensive faulting system. The most superficial one has been overexploited. In the deepest water body, the piezometric levels are below the sea level (with depths in boreholes ranging 350–400 m), whereas the piezometric level in the intermediate one ranges 96–188 m a.s.l. [17]. The hydrochemical facies of the two deeper water bodies are of sulphate-calcium type. The small

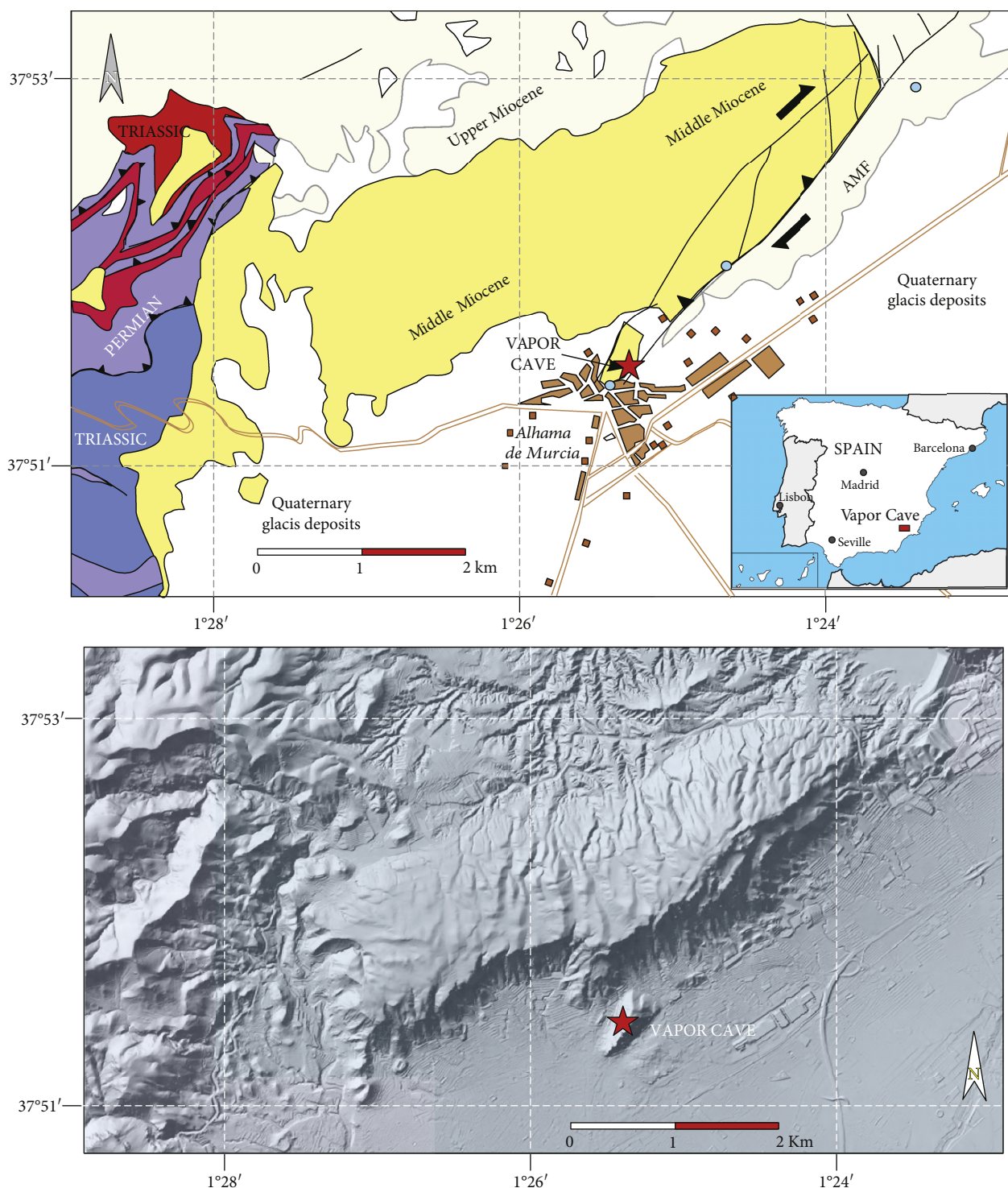


FIGURE 1: Location, local geological settings, and digital terrain model of the study area in relation to the Alhama de Murcia Fault (AMF).

upland where VC is located belongs to the intermediate buckled strata, as well as the thermal boreholes displayed in Figures 2 and 3.

Besides the Alhama de Murcia Fault (NE-SW-trending, Figure 2), the local geometry of the aquifer determines that the VC site is in the middle of consecutive reverse

faults with opposite dips. This faulting system results in an alternation in depth of the black Triassic dolostones (Tr) or Tortonian conglomerates with the younger red sandstones and conglomerates of the middle-lower Tortonian age (A) cross-section of Figure 2 and lithological column of borehole 1 and 2 in Figure 3.

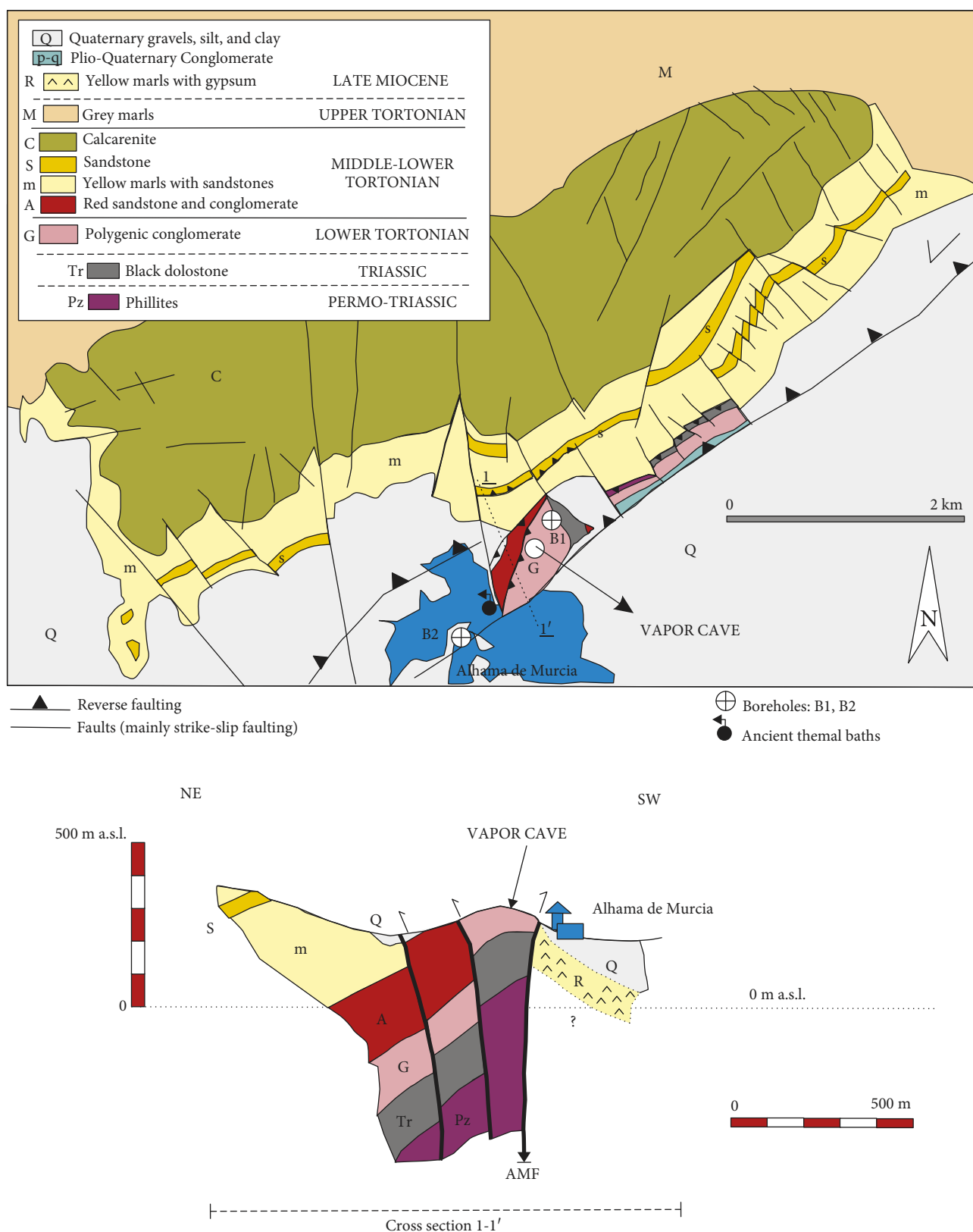


FIGURE 2: Geological map of the study area with main geological units (strata and faults) that set up the local hydrological features in the vicinity of Vapour Cave. Locations of the cave and two key boreholes are indicated and further discussed in Figure 3.



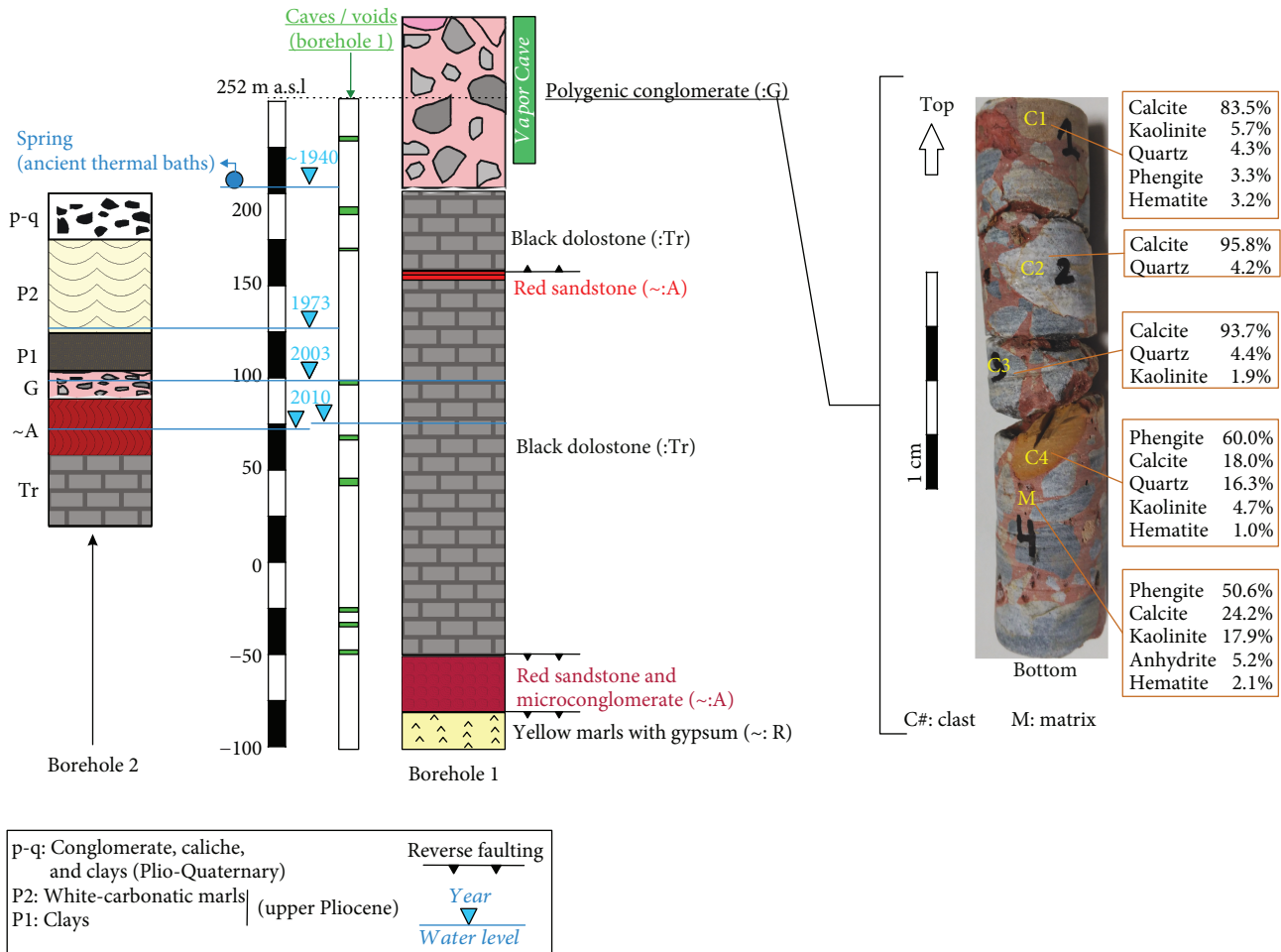


FIGURE 3: Lithological columns of two key boreholes near Vapour Cave; “Cerro del Castillo” (borehole 1) and “Agua de Dios” (borehole 2), both of them linked to the ancient thermal baths of Alhama de Murcia (southern Spain) (adapted from [17] and other unpublished technical reports provided by the Alhama de Murcia city council). Time evolution of water table and depths of the karstified sections (presence of caves and voids) are displayed in relation to lithologic units. The mineral composition (DRX analysis) of both clasts and matrix of a small core sample of the polygenic conglomerate (host rock of Vapour cave) is also detailed.

The Alhama de Murcia Fault would control the flux of groundwater by two ways (Figure 2, low panel with the geological cross section):

- (1) The relative subsidence of the SW block of materials (Quaternary deposits and late Miocene marls with gypsum) provokes the contact of the Triassic dolostones with the impermeable marls, and consequently, the circulation of groundwater towards southeast is hindered.
- (2) The elevation of the northwest block of materials (including the conglomerates hosting Vapour Cave and the lower aquifer of Triassic dolostones) enhanced the circulation of deep and thermal groundwater towards the surface. The local discharge of thermal groundwater was historically located in the layer contact between the dolostones and the overlying conglomerates, just on the natural spring of thermal water that fed the ancient baths at Alhama de Murcia. Nowadays, the piezometric surface is located

at variable depths depending on the extractions destined to irrigation in the whole aquifer.

Figure 3 shows the lithological columns of two key boreholes near VC. Borehole 1 (namely, “Cerro del Castillo”) is 350 m deep, and it is located 197 m far from the entrance to Vapour cave at 252 m a.s.l. Borehole 2 (namely, “Agua de Dios”) is located in the downtown of Alhama de Murcia, and it was drilled up to 180 m depth aimed at supplying hot water to the local thermal baths. This second borehole is at 202 m a.s.l and approximately 780 m far from VC.

Geothermal activity has been reported in both boreholes, with water temperature of 41.4°C at borehole 1 and ranging 39.8–41.0°C at borehole 2 [17, 18], values in concordance with the historical records of temperature for the spring water that fed the ancient thermal baths [19].

Borehole 1 is quite representative of lithology below VC, particularly providing information about the strata just below de Tortonian conglomerates that host the cave. Figure 3 also includes the mineralogical composition of the polygenic conglomerate of Tortonian age. The mineralogical

composition was analysed by powder X-ray diffraction in a Philips PW 1710/00 diffractometer (National Museum of Natural Sciences, Madrid) using  $\text{CuK}\alpha$  radiation with a Ni filter and a setting of 40 kV and 40 mA. Data were collected and interpreted using the X Powder software package. The qualitative search-matching procedure was based on the ICDD-PDF2 database. There is a high percentage of calcite clasts, which were cemented with a phengite matrix. Thick packets of phengite have been described for the Triassic materials of the Alpujárride Complex in southern Sierra Espuña [20], near the study site but at higher elevations. Phengite of the polygenic conglomerates likely came from weathering of these emerged Triassic materials and then transported with clasts through alluvial systems to the coastline, where conglomerates were formed during the lower Tortonian.

Vapour cave (VC), at 295 m a.s.l., only breaks through the Conglomerates of Tortonian age, but its deepest conduits (roughly at  $-80$  m depth) do not reach the black Triassic dolostones, at least through human-size paths. These dolostones should appear in just 13 meters, approximately, below the bottom of the cave and locally house the water table of the aquifer.

During the drilling works of borehole 1, it was also registered in the lithological column the data of depths with caves and voids due to karstification (Figure 3). Borehole 1 crosses through a 4 m high cave in the polygenic conglomerates, which would be located at 65 m depth in the vertical section of Vapour Cave. The remainder of the caves crossed by borehole 1 is embedded into the layer of Triassic black dolostones, with heights ranging 1 to 5 meters [17]. This karstified section of the dolostone strata proves the existence of other trapped air pockets, particularly below Vapour Cave, with a likely presence of endogenous gases that can be potentially released to the exterior atmosphere throughout the faulting system or smaller fissures. Vapour Cave would be a single case of subterranean air pocket that has made its way outwards through the overlying conglomerates.

The records on borehole 2 confirm a continuous decrease in the piezometric level due to overexploitation of the whole aquifer Santa Ychar-Alhama, from the extinct natural spring (roughly at 200 m a.s.l.) to water below 75 m a.s.l. in the last decade (Figure 3). Therefore, the vadose (unsaturated) conditions of the local aquifer have gone prevailing progressively during the last decades and, consequently, the percentage of air reservoirs in the karstified dolostones has increased in the same way. By considering a current hydrostatic level in the local aquifer in a range of 50–75 m a.s.l. (located in the dolostone strata in both boreholes), the estimated thickness of unsaturated rock (polygenic conglomerates plus black dolostones) below VC would reach 140–165 m.

### 3. Cave Settings

VC represents a chasm in a karstic area of active faulting and developed in a favourably fissured carbonate-cemented conglomerate host rock under hypogene speleogenesis by the upwelling of hydrothermal ( $38$ – $43^\circ\text{C}$ , and 100% relative humidity) and  $\text{CO}_2$ -rich fluids, in or from the zone of fluid-

geodynamic influence. In addition to high air temperatures, VC presents other extreme values of some environmental parameters, such as hypoxic conditions (17%  $\text{O}_2$ ),  $\text{CO}_2$  concentrations that exceed 1%, radon ( $^{222}\text{Rn}$ ) activity with values above  $50\text{ kBq/m}^3$ , and a vertical thermal gradient of  $3.2^\circ\text{C}/100\text{ m}$ . All of these conditions have been associated with the combined effects of tectonic activity and hydrothermalism [16]. The current thermal gradient is capable of sustaining free convection of rising  $\text{H}_2\text{O}_{(\text{V})}$  and  $\text{CO}_2$  outgassed from endogenous  $\text{CO}_2$ -rich waters. Consequently, active carbonic acid dissolution still occurs as a hypogenic agent.

The deepest explored part of VC is associated with ancient Roman baths (*Baños de Alhama*), which were exploited during many cultural periods in the village (Roman, Muslim, Medieval, and Modern ages). Geochemical analyses of groundwater at this location have been performed [21]. Groundwater temperature was found to have a mean value of  $41^\circ\text{C}$ ; a pH of 6.8 was documented, and the concentrations of  $\text{Cl}^-$  and  $\text{HCO}_3^-$  suggested the presence of carbonated water. The isotopic signal of dissolved carbon dioxide ranges  $-8\text{‰} < \delta^{13}\text{C}_{\text{CO}_2} < -4\text{‰}$ , in agreement with the thermal degradation of carbonates.

VC is a hypogenic cave that developed in two well-defined sections, subvertical gallery and vertical shafts, with a total explored depth of 84 m below the surface (Figure 2). The first gallery is a subhorizontal tube with an oval section and 50 m of developed marginal outlets, which contain small voids. It appears to be a master passage sloping downward to 30 m below the surface, which developed in carbonate fissures and near-surface fissures running perpendicular to AMF (Figure 2). It constitutes a laterally extensive network with some small “blind-ended” passages, some of which serve as conduits for lateral migration of fluids to the nearest outlet feature (e.g., “ventilador gallery”). The uppermost part of this gallery exhibits the typical morphogenetic features of an outlet conduit, with some cupolas and vertical channels that rise from the ceiling, one of which connects with the exterior through a single, narrow entrance located at the highest point. The cave entrance is oval-shaped, and the axes measure  $0.6 \times 0.75\text{ m}$ . This master passage is connected to a vertical feeding channel (50 m long) that reaches 84 m deep and is related to the AMF fault plane (NE-SW-trending). The chasm becomes narrower as it reaches the lower vadose zone, and it constitutes the main conduit of rising flow of endogenous gases. The bottom of the feeder channel is obscured by the presence of sediment in-fill and breakdown blocks of conglomerate fallen from upper levels; some swallowing or entrenchment forms can also be distinguished.

### 4. Cave Monitoring, Air Sampling Procedures, and Gas Analytical Techniques

A cave air monitoring, gas sampling, and analytical protocol was developed to obtain the key data for understanding the cave-soil-atmosphere system, including temperature, and gaseous composition of the air (molar fractions of  $\text{CO}_2$  and  $\text{CH}_4$  and  $\delta^{13}\text{C}_{\text{CO}_2}$ ,  $\delta^2\text{H}_{\text{CH}_4}$ , and  $\delta^{13}\text{C}_{\text{CH}_4}$  values). Monitoring

was conducted via *in situ* spot measurements, and air samples were analysed in a laboratory setting.

*In situ* and discrete air sampling as well as spot measurements were conducted during intensive surveys lasting for several hours each. The deepest locations were only accessible by very experienced speleologists with self-contained breathing apparatuses due to low oxygen content, extreme temperatures, and hazardous concentrations of other gases. Consequently, *in situ* measurements and air sampling at these deeper locations were only conducted when feasible, and always with the technical assistance of the Mountain Rescue Team (GERA) of the Firefighter's Service of Madrid Autonomous Region.

Overall, eight air-sampling campaigns were conducted from 2015 to 2017 during September of 2015, March and November of 2016, and April and June–September of 2017. The air-sampling campaigns span two hydrogeological cycles, i.e., from September 2015 to August 2016 and from September 2016 to August/September 2017. The two most recent sampling campaigns (August and September of 2017) also included analyses of  $\delta^2\text{H}_{\text{CH}_4}$  in air samples, which were performed within 2 weeks of sample acquisition. Spot sampling of cave air was typically conducted in a predefined network of points that were spatially distributed at several depths: 2, 15, 30, 50, and 80 m below the surface, as well as in randomly distributed locations throughout the cave (Figure 2). The background atmosphere at the exterior was sampled along a transect from the cave entrance until reaching 32 meters far from it. Cave air and exterior atmosphere samples were collected at 1 m above the floor using a portable air compressor (Aquanic s790) running at  $0.4\text{ L}\cdot\text{min}^{-1}$ . Samples for soil gas analyses were collected at several fixed sites located vertically above the cave, and using a 6 mm OD steel tube with grooved sides at each end, and inserted to a depth of 30–50 cm through undisturbed soils to the bedrock-soil interface. Soil air was extracted using a microdiaphragm gas pump (KNF Neuberger, Freiburg, Germany) at  $3.1\text{ L}\cdot\text{min}^{-1}$  at atmospheric pressure. All air samples were collected into 1 L Tedlar bags with a lock valve design specifically to ensure inertness and gas tightness. Air samples from the soil were collected during the same time slot as the sampling of cave air and background atmosphere at the exterior, then all of them were analysed in the laboratory for determining the gas composition (molar fractions of  $\text{CO}_2$  and  $\text{CH}_4$ , and for  $\delta^{13}\text{C}_{\text{CO}_2}$  and  $\delta^{13}\text{C}_{\text{CH}_4}$  values) within 48 hours following the sample collection on field.

Spot measurements of air temperature, relative humidity, air pressure, and  $\text{CO}_2$  concentrations were also taken in the same cave locations as the cave air samples and from the cave exterior using handheld devices (XP100 and XP200, Lufft) with integrated air pressure sensors (measurement range: 800–1100 mbar, accuracy at  $25^\circ\text{C}$  and 1013.25 mbar, max.  $\pm 0.5$  mbar), an external temperature probe (PT100 1/10 DINB probe, with an accuracy of  $\pm[0.03 + 0.002 \cdot \text{measurement}]$ ), and a capacity probe for relative humidity (measurement range: 0–100%, accuracy:  $\pm 3\%$  above 90%). All devices had certified calibrations. Furthermore, a multigas monitor (MX6 iBrid, Industrial Scientific) was used for quantification of  $\text{CO}_2$  and  $\text{O}_2$  concentrations of cave air, including safety

warning levels, and those of other gases potentially present in hypogene environments (e.g.,  $\text{H}_2\text{S}$ , VOCs, and  $\text{H}_2$ ). Further technical details may be found at <http://www.indsci.com/products/multi-gas-detectors/mx6/>.

During the last two field campaigns, the installation of a multichannel system to collect samples from the cave entrance, and at five depths within the cave, 2, 15, 30, 50, and 80 m below the surface, was completed. This system consists of a 100 m long hose with 5 flexible PVC tubes of 25, 50, 75, and 100 m lengths, respectively, such that the end of each tube is located at the aforementioned depths. Cave air was extracted using a microdiaphragm gas pump (KNF Neuberger, Freiburg, Germany) at  $4.5\text{ L}\cdot\text{min}^{-1}$  at 1.5 bar. To minimize the effects of water condensation in the tubes, a laboratory gas drying unit (i.e., a polycarbonate tube with filters, and filled with Drierite desiccant) was installed before filling the Tedlar bags. Finally, a multigas MX6 iBrid monitor was connected in sequence to the sampling tubes and enclosed in an air-tight box to measure the gas concentrations of the air collected at each depth.

Air samples were analysed at the National Museum of Natural Sciences (Spanish National Research Council-CSIC) for determining  $\text{CO}_2$  and  $\text{CH}_4$  molar fractions, as well as the isotopic  $\delta^{13}\text{C}$  values for both gases using wavelength-scanned cavity ring-down spectroscopy (CRDS-WS). A Picarro G2201-i CRDS analyser (Picarro Inc., USA) was used to quantify the isotopologues of carbon dioxide and methane and to automatically calculate the carbon isotopic value for both gases with high precision. According to the manufacturer's technical specifications, this CRDS analyser measures the isotopologues of carbon dioxide ( $^{12}\text{CO}_2$  and  $^{13}\text{CO}_2$ ) with a precision of 200 ppb ( $\pm 0.05$  of reading) and 10 ppb ( $\pm 0.05$  of reading) for  $^{12}\text{CO}_2$  and  $^{13}\text{CO}_2$ , respectively, resulting in a precision greater than 0.16‰ for  $\delta^{13}\text{C}_{\text{CO}_2}$  after 5 min of analysis. The measurements of methane isotopologues ( $^{12}\text{CH}_4$  and  $^{13}\text{CH}_4$ ) reached a precision of 5 ppb ( $\pm 0.05$  of reading) and 1 ppb ( $\pm 0.05$  of reading) for  $^{12}\text{CH}_4$  and  $^{13}\text{CH}_4$ , respectively, resulting in a precision greater than 1.15‰ for  $\delta^{13}\text{C}_{\text{CH}_4}$  after 5 min of analysis.

Air samples in duplicate Tedlar bags were sent to the Institute for Marine and Atmospheric research (IMAU, Utrecht University) for high-precision measurements of hydrogen isotopes on atmospheric methane (and  $\delta^{13}\text{C}_{\text{CH}_4}$  for some samples) by using continuous-flow isotope ratio mass spectrometry (CF-IRMS) and following previously published methods [22]. This method separates  $\text{CH}_4$  from other air components by utilizing purely physical processes based on temperature, time, and mechanical valve switching (i.e., without any added chemicals), and the purified sample is then pyrolysed to  $\text{H}_2$  for stable isotope measurements. This analytical procedure allows high-precision measurements of  $\delta\text{D}$  and  $\delta^{13}\text{C}$  from atmospheric  $\text{CH}_4$  samples, with typical reproducibility of  $\pm 0.07\text{‰}$  for  $\delta^{13}\text{C}$ , 2.3‰ for  $\delta\text{D}$ , and 17 ppb for  $\text{CH}_4$  concentrations. For the CF-IRMS analyses, the amount of sample gas was adjusted to yield the same amount of  $\text{CH}_4$  for each measurement. The general atmospheric measurement setup of the CF-IRMS analyser was adjusted by adding an ascarite/Mg ( $\text{ClO}_4$ )<sub>2</sub> filter to remove the preexisting large amounts of  $\text{CO}_2$ . To measure samples

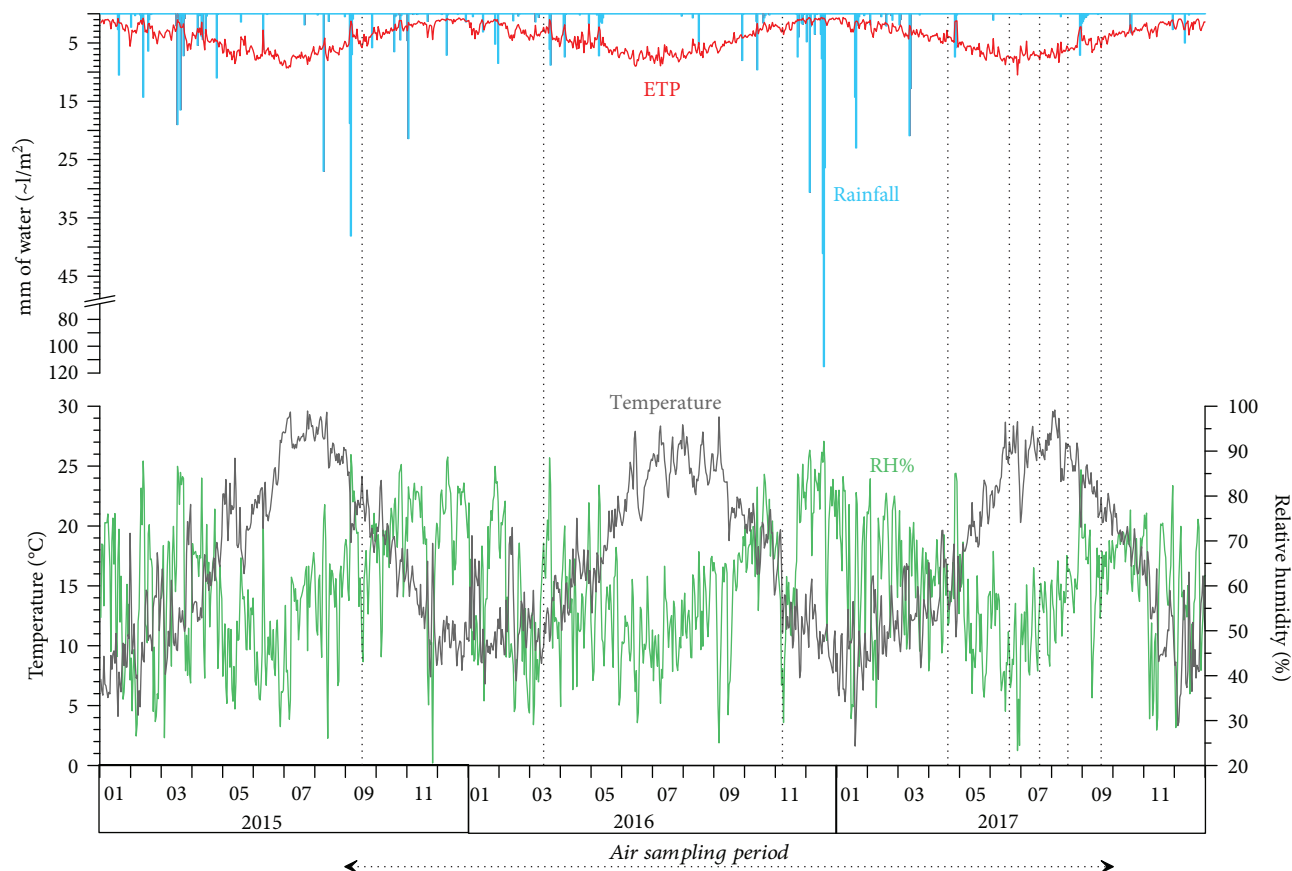


FIGURE 4: Daily meteorological conditions in the study area from January 2015 to December 2017, including the dates of the air sampling campaigns (dotted vertical lines). Parameters; air temperature, relative humidity, rainfall, and evapotranspiration according to the Penman-Monteith FAO parameterization scheme [25].

with lower  $\text{CH}_4$  content, this method was further adjusted to extract  $\text{CH}_4$  from larger volumes of air. The stable carbon and hydrogen isotope compositions of both gases ( $\text{CO}_2$  and  $\text{CH}_4$ ) are expressed as  $\delta^{13}\text{C}$  and  $\delta^2\text{H}$  relative to standards Vienna Pee Dee Belemnite (VPDB) and Vienna Standard Mean Ocean Water (VSMOW).

Three in-house standards with certified gas mixtures and known  $\text{CO}_2$  and  $\text{CH}_4$  concentrations (6993 ppm, 399 ppm, and zero- $\text{CO}_2$  and 0.5 ppm, 1.7 ppm, and zero- $\text{CH}_4$ , respectively, supplied by Praxair Spain) were processed regularly at the beginning and at the end of each analytical session to verify the proper functioning of the CRDS analyser. Further details about the methodological procedures and quality results can be found elsewhere [23]. Additionally, we periodically evaluated  $\delta^{13}\text{C}_{\text{CH}_4}$  measurements by processing diluted air samples extracted from the following standard gases with certified methane stable isotope ratios: T-iso3 (250 ppm  $\text{CH}_4$  and  $-38.3\text{‰}$   $\delta^{13}\text{C}_{\text{CH}_4}$ ) and L-iso1 (2500 ppm  $\text{CH}_4$  and  $-66.5\text{‰}$   $\delta^{13}\text{C}_{\text{CH}_4}$ ), both supplied by Isometrics Instruments (Canada). In-house standards were also subjected to quality control by comparing the results obtained with the Picarro G2201-i analyser with duplicated bags collected from cylinders and subsequently analysed independently in the greenhouse gas laboratory at the Royal Holloway University of London (RHUL). There, gas concentrations were analysed with a Picarro G1301 CRDS analyser, and  $\delta^{13}\text{C}_{\text{CO}_2}$  and

$\delta^{13}\text{C}_{\text{CH}_4}$  were measured in triplicate by CF GC-IRMS using a GV Instruments Trace Gas e IsoPrime system [24]. Finally, duplicated air samples collected in situ were also analysed for  $\delta^{13}\text{C}_{\text{CH}_4}$  by CF-IRMS (IMAU, Utrecht) and then compared with measurements provided by the Picarro G2201 CRDS analyser, confirming an agreement between both analyses. Overall, these internal and intercomparison procedures periodically validated that the performance specifications regarding  $\text{CO}_2$  and  $\text{CH}_4$  analyses via a CRDS analyser were met.

## 5. Results

During the first hydrological cycle, the timing of access to the cave and air sampling was determined by availability of the qualified speleologists abovementioned. The sampling campaigns were intensified during the second hydrological cycle thanks to the installation of the multichannel system to collect samples from the cave entrance, i.e., without need to access the cave. In this period of two hydrological cycles, 8 field campaigns were conducted, 5 of which included sampling up to 30 m deep, and 3 of which included air sampling up to 80 m deep.

The daily meteorological conditions in the study area during the sampling period are shown in Figure 4, including the dates of the air sampling campaigns. Table 1 summarizes



TABLE 1: Climate conditions in the study are during the last two decades, including the annual-average values of main climatic parameters: air temperature, relative humidity, rainfall, and evapotranspiration according to the Penman-Monteith FAO parameterization scheme [25].

Hydrological year	T_avgr (°C)	RH_avgr (%)	Rainfall (mm) ETP	PM_FAO (mm)
Sept. '96–Aug. '97	17.08	68.7	271	1210
Sept. '97–Aug. '98	17.70	67.1	242	1239
Sept. '98–Aug. '99	17.35	64.1	206	1268
Sept. '99–Aug. '00	17.36	65.5	135	1250
Sept. '00–Aug. '01	17.95	63.3	183	1319
Sept. '01–Aug. '02	17.20	66.6	416	1192
Sept. '02–Aug. '03	18.09	62.7	212	1262
Sept. '03–Aug. '04	17.55	67.7	360	1161
Sept. '04–Aug. '05	17.22	67.2	132	1340
Sept. '05–Aug. '06	17.19	70.2	316	1308
Sept. '06–Aug. '07	17.70	69.2	344	1359
Sept. '07–Aug. '08	16.81	67.1	305	1376
Sept. '08–Aug. '09	16.54	59.8	319	1423
Sept. '09–Aug. '10	16.64	66.2	520	1330
Sept. '10–Aug. '11	16.59	64.1	189	1384
Sept. '11–Aug. '12	17.06	58.3	128	1575
Sept. '12–Aug. '13	16.52	59.9	355	1424
Sept. '13–Aug. '14	17.41	57.4	95	1560
Sept. '14–Aug. '15	17.60	58.6	266	1531
Sept. '15–Aug. '16	17.32	59.5	187	1491
Sept. '16–Aug. '17	17.51	60.8	382	1459
Average year	17.26	64.0	265	1355

the evolution of climate during the last two decades, including the annual-average values of main climatic parameters: air temperature, relative humidity, rainfall, and evapotranspiration. These climate data sets were supplied by a meteorological station at Alhama de Murcia, belonging to the Network of the Agricultural Information System of Murcia (<http://siam.imida.es/>) and located at 169 m a.s.l and less than 7 km far from Vapour Cave.

The area has a semiarid Mediterranean climate. The average annual temperature is 17.26°C, ranging 16.5–18.1°C over the last two decades. During the sampling period, the minimum values of the daily average temperature were recorded in the winter months of December/January, ranging 8.5–10.4°C, and the highs in the summer months of July/August (25.7–27.8°C). The average annual relative humidity (RH) is 64%, historically ranging from 57.4% to 70.2%. During the sampling period, the minimum values of the daily average RH were recorded from May to July, usually below 50%, and the highs in October (>60%).

Rainfall in the area is typical of a semiarid climate: 265 mm/year on average during the last two decades, with some cycles under very dry conditions (<100 mm/year). Annual rainfall for a whole hydrological cycle has ranged from 187 to 380 mm during the sampling period (2015–2017). The annual evapotranspiration is five times higher than precipitations (this difference factor ranged from 3.8 to 7.9 during the last two hydrological cycles), which results on prevailing xeric conditions.

It has a very marked dry season in the summer, a period in which more than 3 months (May to July/August) are usually registered without any precipitation or, at least, rainfall lower than 5 mm/month. During the rest of the year, precipitation events are very scarce and are distributed between winter, spring, and autumn, but without a constant pattern (e.g., very low rainfall is registered during some winter months). Low rainfall on an annual scale is opposed to the existence of some torrential events, for instance, 194 mm during just five days (15–19 December 2016).

The mean value of measured CO<sub>2</sub> and CH<sub>4</sub> concentrations and their stable isotopic compositions ( $\delta^{13}\text{C}_{\text{CO}_2}$  and  $\delta^{13}\text{C}_{\text{CH}_4}$ ) in the cave environment, above-cave soils, and local exterior atmosphere are shown in Table 2. Soil-CO<sub>2</sub> concentrations were relatively low, with heavy  $\delta^{13}\text{C}_{\text{CO}_2}$  values in all samples, and range from 527 ppm in September of 2015 to 2034 ppm in September of 2017 following moderate rainfall (3.5 L m<sup>-2</sup>). Values of  $\delta^{13}\text{C}$ -CO<sub>2</sub> of soil air ranging from -9.8 to -15.9‰ are consistent with a local semiarid climate.

The local outdoor atmosphere has mean values of CO<sub>2</sub> and CH<sub>4</sub> concentrations slightly above the recent global monthly mean CO<sub>2</sub> (roughly 405 ppm and 1.85 ppm, respectively, check <https://www.esrl.noaa.gov/gmd/ccgg/data-products.html> for our monitoring period). The highest CO<sub>2</sub> values were recorded in the autumn, coinciding with colder outdoor air temperatures.

TABLE 2: Mean measurements (and  $\pm$ sd: standard deviation) for the concentrations of  $\text{CO}_2$  and  $\text{CH}_4$ , and their stable isotopic compositions ( $\delta^{13}\text{C}_{\text{CO}_2}$  and  $\delta^{13}\text{C}_{\text{CH}_4}$ ) in the cave environment, and in vertically adjacent soils and the local outdoor atmosphere.

	$\text{CO}_2$ (ppm)		$\delta^{13}\text{C}_{\text{CO}_2}$ (%)		$\text{CH}_4$ (ppm)		$\delta^{13}\text{C}_{\text{CH}_4}$ (%)		$n$
	avgr	$\pm$ sd	avgr	$\pm$ sd	avgr	$\pm$ sd	avgr	$\pm$ sd	
Sep. '15									
Soil	527	6	-9.83	0.15	1.80	0.00	-46.89	0.80	4
Cave	8257	1795	-5.77	0.56	1.09	0.16	-42.57	3.31	8
Ext.	411	2.06	-8.26	0.16	1.91	0.01	-47.55	0.39	4
Mar. '16									
Soil	810	218	-11.80	0.53	1.86	0.12	-46.36	0.57	4
Cave	9040	2575	-6.29	0.83	2.62	0.49	-49.17	1.95	9
Ext.	—	—	—	—	2.06	0.02	-48.10	0.45	4
Nov. '16									
Soil	1907	929	-15.60	2.33	0.52	0.55	-44.07	5.30	5
Cave	11.964	1535	-5.83	0.32	0.87	0.06	-44.43	16.42	9
Ext.	503	153	-7.84	0.58	2.03	0.01	—	4.28	4
Apr. '17									
Soil	1472	1430	-15.87	0.56	0.71	0.61	-42.66	6.34	3
Cave	7507	2036	-6.35	0.43	1.08	0.14	-45.95	1.64	10
Ext.	465	20	-10.71	0.51	2.01	0.01	-51.93	0.64	3
Jun. '17									
Soil	1035	702	-12.46	1.60	1.76	0.37	-45.33	0.20	3
Cave	7775	979	-5.56	0.12	0.95	0.06	-35.29	2.22	5
Ext.	434	271	-9.25	1.29	2.16	0.08	—	5.49	4
Jul. '17									
Soil	639	87	-12.79	0.57	1.88	0.18	-47.20	1.72	3
Cave	8998	1718	-5.72	0.08	0.94	0.22	-41.40	16.31	3
Ext.	433	193	-8.08	1.10	2.04	0.04	—	0.83	3
Aug. '17									
Soil	649	329	-11.07	0.18	1.94	0.12	-47.68	3.39	4
Cave	10.441	1230	-5.35	0.16	0.89	0.12	-38.66	5.31	6
Ext.	434	16	-9.07	0.65	2.00	0.00	-53.08	0.46	3
Sep. '17									
Soil	2034	646	-13.45	1.89	0.50	0.06	-62.89	4.02	3
Cave	12.251	3251	-5.60	0.50	1.07	0.09	-39.41	7.09	8
Ext.	417	9	-8.65	0.09	1.99	0.00	—	1.23	3
Average									
Soil	<b>1134</b>	<b>543</b>	<b>-12.86</b>	<b>0.98</b>	<b>1.37</b>	<b>0.25</b>	<b>-47.88</b>	<b>2.79</b>	<b>29</b>
Cave	<b>9529</b>	<b>1890</b>	<b>-5.81</b>	<b>0.38</b>	<b>1.19</b>	<b>0.17</b>	<b>-42.11</b>	<b>6.78</b>	<b>58</b>
Ext.	<b>442</b>	<b>95</b>	<b>-8.84</b>	<b>0.63</b>	<b>2.02</b>	<b>0.02</b>	<b>-50.17</b>	<b>1.72</b>	<b>28</b>

All of the underground samples presented higher concentrations of  $\text{CO}_2$  and higher  $\delta^{13}\text{C}_{\text{CO}_2}$  than those of the exterior atmosphere and soil air. The underground air exhibits high concentrations of  $\text{CO}_2$  that generally increase with depth. This general trend is broken at 15 m depth by the existence of an external air intake through the “Ventilador” gallery (Figure 5) that causes a decrease in the concentration of  $\text{CO}_2$  and a concomitant increase in the concentration of  $\text{CH}_4$ . The effect of this air intake intensifies during cold periods (e.g., November of 2016), and it significantly influences the thermal and  $\text{O}_2$  profiles. The mean annual

concentration of  $\text{CH}_4$  in the underground air is lower than that of both the soil and the outdoor atmosphere. Only in March of 2016 were  $\text{CH}_4$  concentrations above the atmospheric background (ranging 2.3–3.4 ppm) detected. The  $\delta^{13}\text{C}_{\text{CO}_2}$  values of underground air ranged from -4.5 to -7.5‰, and the  $\text{CH}_4$  molar fractions and both  $\delta\text{D}$  and  $\delta^{13}\text{C}$  values ranged from -77 to 48‰ and -52 to -30‰, respectively.

The temperature profile of the cave interior reveals three clear divisions (Figure 5): (1) a shallow thermal zone from 0 to 15 m depth controlled by exterior air influx, (2) a

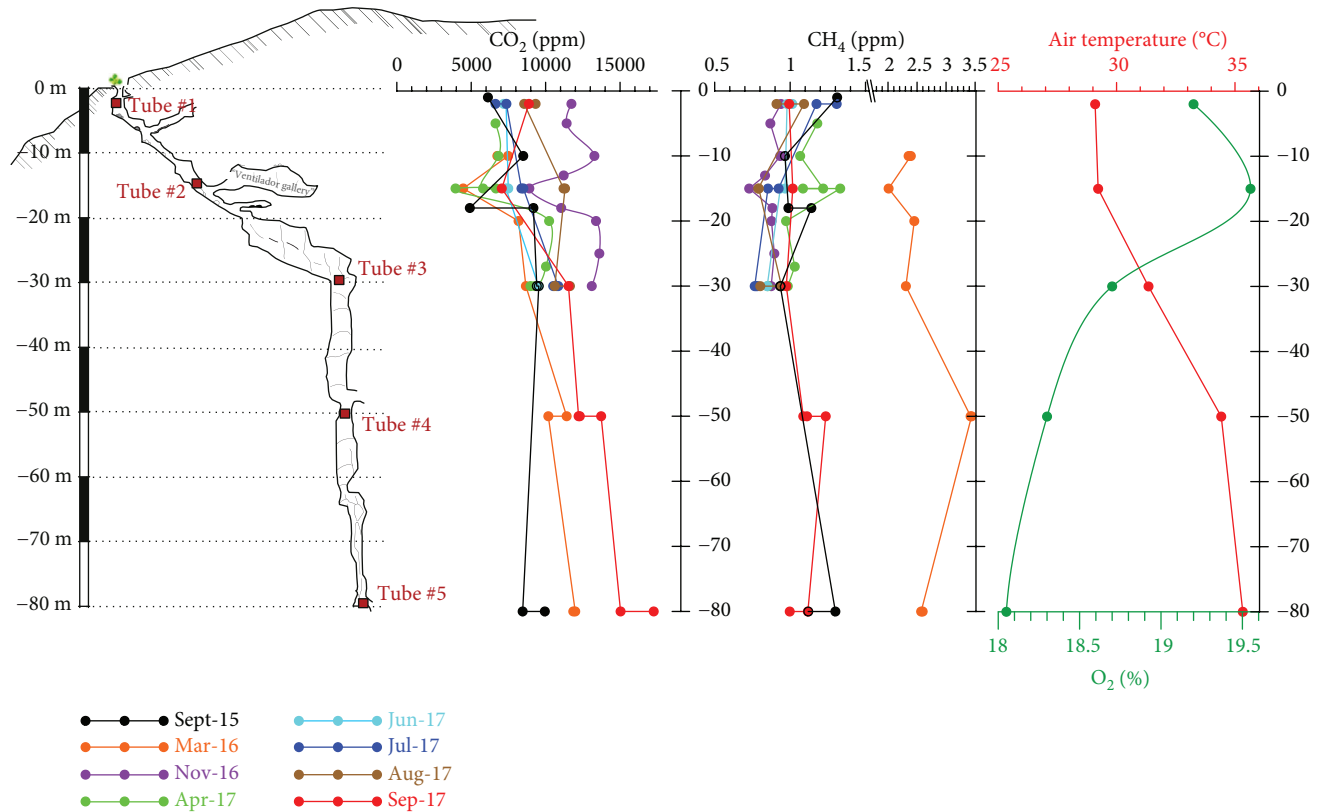


FIGURE 5: Spatiotemporal evolution of temperature and gaseous composition of cave air ( $\text{CO}_2$ ,  $\text{CH}_4$ , and  $\text{O}_2$  contents) along a vertical profile.

heterothermal medium zone between 15 and 50 m with a marked increase in temperature by  $5^\circ\text{C}$  at 35 m depth, and (3) the thermal deep zone with a progressive increase in temperature controlled by the geothermal gradient.

## 6. Discussion

**6.1. Geochemical Tracing of  $\text{CO}_2$  Sources and Dynamic at VC.** The active hypogene speleogenesis at VC is mainly controlled by the upwelling airflow from the zone of fluid-geodynamic influence associated with an active fault with frequent microseisms [26]. Cave air samples have a remarkably high concentration of  $\text{CO}_2$  with heavy  $\delta^{13}\text{C}_{\text{CO}_2}$  values ( $-5.81\text{‰}$  on average, and ranging from  $-7.40$  to  $-4.67\text{‰}$ ), which indicates a clear deep endogenous source of  $\text{CO}_2$  in cave air. These  $\delta^{13}\text{C}_{\text{CO}_2}$  values in air are also in agreement with those measured in  $\text{CO}_2$ -rich thermal waters of the aquifer spatially associated with the active fault (between  $-8.1$  and  $-3.8\text{‰}$  [21]). Therefore, degassing from  $\text{CO}_2$ -rich groundwater and deep-sourcing geothermal  $\text{CO}_2$  seem to be the prevailing processes responsible for the high abundance of  $\text{CO}_2$ , and its heavier carbon isotopic composition. According to the historical piezometric records from some near boreholes, the local water level matches the layer of Triassic black dolostones below VC (roughly 140–165 m), which are highly karstified due to the chemical aggressiveness of this  $\text{CO}_2$ -rich groundwater. The network of karstic voids/caves and fissures below VC would favour the diffusion and convection of

deep-endogenous gases to the upper layers of the aquifer, including the polygenic conglomerates hosting VC.

Similar carbon isotopic ratios have been described for soil air samples from hydrothermal areas within wider volcanic regions [27] and magma-derived  $\text{CO}_2$  emissions [28]. Other studies at hydrothermal sites have described wider ranges of carbon isotope composition of  $\text{CO}_2$  (e.g., from  $-2.4$  to  $-7.8\text{‰}$  in submarine hydrothermal vents [29] and from  $-1.0$  to  $-9.1\text{‰}$  in hot springs [30]). Taking these  $\delta^{13}\text{C}_{\text{CO}_2}$  values as references, a higher  $\delta^{13}\text{C}_{\text{CO}_2}$  may indicate the addition of  $\text{CO}_2$  directly from volcanic sources [31] or from underlying sedimentary rocks containing more marine carbonate minerals (i.e.,  $\text{CO}_2$  produced mainly by thermal decarbonation [32]). On the contrary, lighter  $\delta^{13}\text{C}_{\text{CO}_2}$  values suggest a likely contamination by crustal organic sediments [33].

A Keeling analysis and modelling of the stable isotope fractionation of  $\text{CO}_2$  and  $\text{CH}_4$  was used to identify and assess the processes as consumption, accumulation, and mobilization (e.g., bacterial oxidation of  $\text{CH}_4$  or diffusion of soil-derived or deep-sourced  $\text{CO}_2$ ), as mixture of gases with distinguishable origins, or resulting from different consumption or production processes. In the case of  $\text{CO}_2$  analysis for VC (Figure 6), the Keeling diagram incorporates the assumption that each data point corresponds to the gas composition of cave air, including molar fraction and isotopic value. It represents a mixture of two end-member gases: local atmosphere and pure  $\text{CO}_2$  that have been added to the cave air to produce the composition at the observation point. The

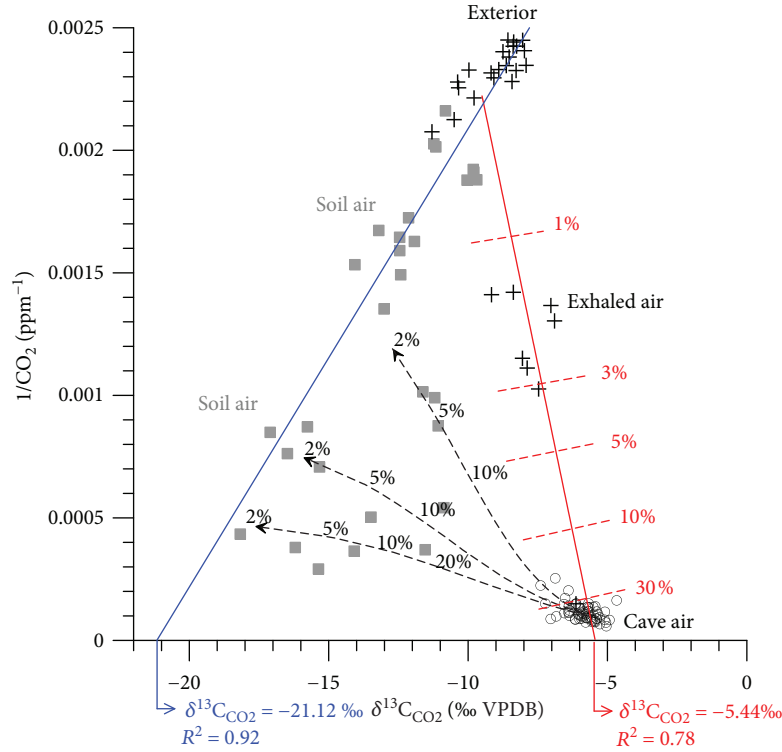


FIGURE 6: Plot of  $1/\text{CO}_2$  versus  $\delta^{13}\text{C}_{\text{CO}_2}$  for soil (grey squares) and cave air (open circles). The composition of the local atmosphere is indicated by crosshairs, distinguishing those samples of exhaled air collected just inside the cave entrance. The Keeling functions fit the set of data points representing a two-component mixture of average atmospheric air with an additional source of  $\text{CO}_2$ , either the pure component from the soil (blue straight line, without considering the dispersed data point towards cave air), or from deep endogenous air (red straight line). Extrapolation down to the X-axis yields the  $\delta^{13}\text{C}_{\text{CO}_2}$  of the  $\text{CO}_2$  source: soil-derived  $\text{CO}_2$  ( $-21.12\text{‰}$ ,  $R^2 = 0.92$ ) and deep endogenous  $\text{CO}_2$  ( $-5.44\text{‰}$ ,  $R^2 = 0.78$ ).

Keeling plot reveals the isotopic composition of this pure  $\text{CO}_2$  by extrapolating the straight line joining the atmospheric end-member to the data point under consideration, as far as its intersection with the  $\delta^{13}\text{C}_{\text{CO}_2}$  axis [34]. This  $\delta^{13}\text{C}_{\text{CO}_2}$  value is only an apparent composition and does not correspond to a real source of  $\text{CO}_2$  if there are other processes occurring (e.g., diffusion or mixing with several pure  $\text{CO}_2$  sources that have more than a single composition for  $\delta^{13}\text{C}_{\text{CO}_2}$ ).

Thus, two sources of  $\text{CO}_2$  are considered in this analysis through Keeling diagrams, either from soil or from deep endogenous air. The red keeling function in Figure 6 shows the effects of mixing between the atmosphere, and the composition of pure deep endogenous  $\text{CO}_2$ , considering a composition close to the maximum  $\text{CO}_2$  concentration and  $\delta^{13}\text{C}_{\text{CO}_2}$  values measured for the cave air samples (17,623 ppm and  $-4.67\text{‰}$ , respectively) and the average composition of the local atmosphere (442 ppm and  $-9.01\text{‰}$ ). Red dashed lines are contours of equal mixing ratios labelled as % of pure  $\text{CO}_2$  remaining in the cave air samples, including the exhaled air from the cave entrance. According to this model, the underground air at VC usually maintains more than 30% pure theoretical  $\text{CO}_2$  (with a  $\delta^{13}\text{C}_{\text{CO}_2} = -5.44\text{‰}$ ) added from a deep endogenous source. The mixing process with the local atmosphere increases as the upwelling flux of air travels to the cave entrance; thus, the exhaled air to the outdoor atmosphere represents between 1 and 3% of this pure theoretical  $\text{CO}_2$  added from a deep endogenous source.

The blue Keeling function in Figure 6 shows the effects of mixing between the atmosphere and the composition of the theoretical pure soil  $\text{CO}_2$ . The soil-derived  $\text{CO}_2$  includes all  $\text{CO}_2$  originally generated within the soil (i.e., from root respiration and soil organic matter degradation) as well as some subsequent processes (e.g., direct gas diffusion mainly from deeper soil layers or previously accumulated in the fissures, fractures, and pore spaces of rocks in the vadose zone). The extrapolation down to the X-axis gives  $\delta^{13}\text{C}_{\text{CO}_2}$  of the soil-derived  $\text{CO}_2$  source of  $-21.12\text{‰}$  ( $R^2 = 0.92$ ). This value is consistent with  $\text{CO}_2$  derived from the decomposition of C3 biomass ( $-27 \pm 3\text{‰}$ ) plus a 4.4‰ diffusional enrichment [35], but clearly distinguishable from the epigenetic caves characterized by a lighter end-member for soil  $\text{CO}_2$  production with  $\delta^{13}\text{C}_{\text{CO}_2} = -26\text{‰}$  or less [36].

This  $\delta^{13}\text{C}_{\text{CO}_2}$  value also indicates that  $\text{CO}_2$  produced by microbial respiration in soils containing organic material from C3 vegetation might not be the only process responsible for the concentration and carbon isotopic composition of soil  $\text{CO}_2$ . In fact, there are many data points for soil air that scatter closely under the best-fit line (blue Keeling plot in Figure 4), and therefore, these points seem to represent mixtures between atmospheric air and more than one single pure  $\text{CO}_2$  end-member (i.e., they are not only from soil-derived  $\text{CO}_2$ ).

Because the scatter of soil points is high, we have established a complementary model to explain the outliers of soil gas composition. This is based on how upwelling flow of deep



endogenous gases in this cave influences the  $\delta^{13}\text{C}_{\text{CO}_2}$  values of the soil air, which is likely due to an intense  $\text{CO}_2$  diffusion from the cave to the soil layers located immediately above. This effect is noticeable because the data points from the soil air with the highest  $\text{CO}_2$  concentrations tend to drift away in a perpendicular direction from the mixing line exterior-soil (blue keeling function in Figure 3) towards the data set of cave air with higher  $\text{CO}_2$  content and heavier  $\delta^{13}\text{C}_{\text{CO}_2}$ .

The curved dashed arrows in Figure 6 show the kinetic fractionation trajectory of deep endogenous  $\text{CO}_2$  due to its upwards diffusion from cave air to soil layers through fissures and small-size cracks of deep soil-epikarst. Gas diffusion, driven by concentration gradients according to Fick's law, may produce  $^{13}\text{C}$  depletion in the diffusing gas collected in the soil air samples, and consequently, the residual  $\text{CO}_2$  gas in cave air will be  $^{13}\text{C}$ -enriched. In any case, the diffusing gas that reaches the soil environment is  $^{13}\text{C}$ -enriched with respect to the soil-derived  $\text{CO}_2$  (i.e., from root respiration and soil organic matter degradation), which is identified with the data pairs better aligned to the mixing line exterior soil (blue Keeling plot in Figure 6). Gas diffusion is modelled by means of a Rayleigh-type distillation process with several kinetic fractionation coefficients (3.05‰, 2.60‰, and 1.77‰) in the function of the  $\text{CO}_2$  gradient between cave air and the deepest layers of soil. The Rayleigh equation is an exponential relation that describes the partitioning of isotopes between two reservoirs as one reservoir decreases in size, in this case the  $\text{CO}_2$  content in soil air. These kinetic fractionation coefficients result from fitting the Rayleigh-type distillation curves considering the average values of  $\text{CO}_2$  concentration and  $\delta^{13}\text{C}_{\text{CO}_2}$  of cave air (9529 ppm and  $-5.81\text{‰}$ , respectively) and three representative soil  $\text{CO}_2$  values ([2000 ppm, 18.48‰], [1175 ppm, 16.62‰], and [600 ppm, 13.18‰]), in accordance to the mixing line between soil air and local atmosphere (blue keeling function in Figure 6). There is an intense vapour condensation on cave wall and ceilings during the upwards flux of warm and humid air, which hinders the gaseous connection between the cave environment and the above soil layer through fissures, small size cracks, and the connected porous system. As a consequence, the kinetic fractionation coefficients used to model the  $\text{CO}_2$  diffusion are lower than the theoretical mass-dependent fractionation between  $^{12}\text{CO}_2$  and  $^{13}\text{CO}_2$  during diffusion within the external soil layers (4.4‰ [37]). Each diffusion curve has been labelled as percentage of deep endogenous  $\text{CO}_2$  that remains in the soil air after the gas diffusion process occurring between cave and soil. Some soil samples showed a remaining deep endogenous  $\text{CO}_2$  that ranges between 5% and 10%, which demonstrates that the upwelling flow of geogenic  $\text{CO}_2$  has a clear influence on the external soil above the cave.

**6.2. Sources and Sink Processes during Migration and Upwelling of Deep Endogenous Methane.** The  $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$  variations in cave air provide considerable insight into the nature of gas exchange and consumption processes controlling the  $\text{CH}_4$  dynamic in underground environments. A key reference point in the data interpretation is that the background atmosphere usually has around 1.8 ppm of

$\text{CH}_4$  and its carbon and hydrogen isotopic composition ( $\delta^{13}\text{C}_{\text{CH}_4} \approx -47\text{‰}$  VPDB,  $\delta^2\text{H}_{\text{CH}_4} \approx -100\text{‰}$  VSMOW) is a product of inputs from an isotopically wide range of sources. The  $\text{CH}_4$  concentration of cave air in epigenetic caves and, in general, in well-ventilated caves independently of their speleogenesis mechanisms are often depleted, confirming that subterranean environments may represent an overlooked sink for atmospheric  $\text{CH}_4$  [23, 38–44] and, further, it is rapidly consumed in caves on time scales ranging from hours to days [23, 39]. On the opposite case, underground air of some hypogene caves may contain unusually high levels of methane (up to 3%, e.g., Movile Cave) related to the action of chemoautotrophic bacteria [45], and others have moderate  $\text{CH}_4$  concentrations, just above the atmospheric background, related to  $\text{CH}_4$  outgassing from spring water in sulphuric acid hypogenic caves (e.g., <4 ppm  $\text{CH}_4$  at Cueva Villa Luz [7]).

The variations of  $\delta^{13}\text{C}_{\text{CH}_4}$  as a function of methane concentrations in air of VC are illustrated in Figure 7, compared to a standard composition of the local atmosphere ( $\text{CH}_4$ : 2.02 ppm and  $\delta^{13}\text{C}_{\text{CH}_4}$ : 50.17‰). The most relevant fact is that some noticeable concentrations of deep endogenous methane have been occasionally registered. Thus, in the deeper sites of this chasm (below 30 m and, particularly, at 50 m depth), deep endogenous  $\text{CH}_4$  reaches values higher than the atmospheric background (ranging 2.3 to 3.4 ppm) with  $\delta^{13}\text{C}_{\text{CH}_4}$  values, a bit lighter with respect to those found in the local atmosphere (Figure 7). These data were registered for the first time during March of 2016, but no more evidences of high  $\text{CH}_4$  concentration have been observed in the subsequent surveys on field for air sampling. These high concentrations of deep endogenous  $\text{CH}_4$  denote a more intense migration of endogenous fluids through the upper vadose zone, which could be related with an increase in regional seismotectonic activity.

The general trend of the scattered data points in Figure 7 is that smaller  $\text{CH}_4$  concentrations of cave air are associated with the most  $^{13}\text{C}$ -enriched  $\text{CH}_4$ . These data suggest that methanotrophic oxidizing bacteria (MOB) seem to be the main responsible for consumption  $\text{CH}_4$  in cave air. Curves of Figure 7 fit the locus compositions formed by MOB consumption of atmospheric  $\text{CH}_4$ , modelled as a Rayleigh process using several kinetic fractionation factors (F) and considering as starting point the maximum  $\text{CH}_4$  concentration registered at 50 m depth in March of 2016, as a clear example of a deep endogenous source of this gas. This model for  $\text{CH}_4$  consumption by MOB seems to work, since cave air samples with  $\delta^{13}\text{C}_{\text{CH}_4}$  values heavier than  $-50\text{‰}$  are located within the plotted area defined by these distillation curves, and, likewise, data pairs with heavier  $\delta^{13}\text{C}_{\text{CH}_4}$  match with distillation curves with higher fractionation factors.

As a reference, the magnitude of kinetic fractionation factors associated with methane oxidation varies between 1.009 in anoxic aqueous environments [46] and 1.025–1.049 during gas transport in soils above landfill. The fractionation factors for samples of VC are within the range calculated for the aerobic oxidation of  $\text{CH}_4$  from laboratory cultures of methanotrophs ([47, 48] and references therein) and field studies on the vadose zone above

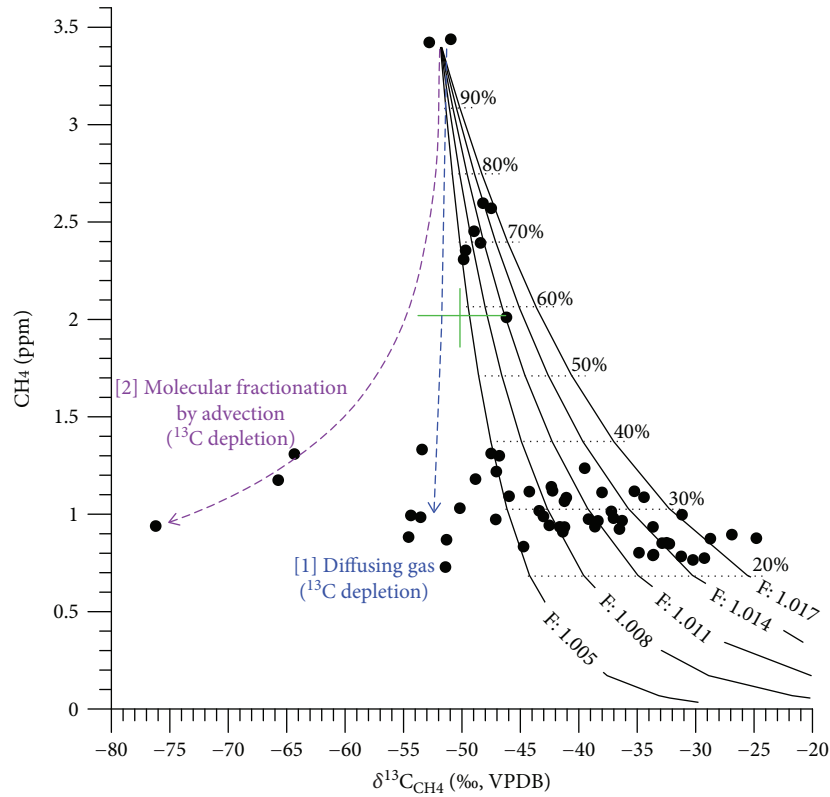


FIGURE 7: Relationship between  $\text{CH}_4$  and  $\delta^{13}\text{C}-\text{CH}_4$  in underground air of Vapour Cave (VC; closed circles). The standard composition of the local background atmosphere is indicated by the green crosshairs (as a reference, the  $\delta^{13}\text{C}_{\text{CH}_4}$  value of this crosshairs separates the  $\text{CH}_4$  involved in bacterial oxidation from  $\text{CH}_4$  coming from other sources). Continuous curves show the locus of compositions formed by methanotrophic consumption of atmospheric methane modelled as a Rayleigh process using several kinetic fractionation factors ( $F$ : from 1.005 to 1.017). Horizontal dotted lines with labels show the percentage of deep endogenous methane remaining in cave air after methanotrophic consumption. Curves for isotopic fractionation by diffusion and molecular fractionation by advection are only inferred, but not modelled.

methanogenic aquifers [49]. However, some curves are defined by fractionation factors smaller than those determined in situ in soils [50–52].

Labels of the distillation curves in Figure 7 show the percentage of deep endogenous  $\text{CH}_4$  remaining in cave air after its consumption by MOB. In general, cave air samples with subatmospheric  $\text{CH}_4$  (<1.3 ppm and  $\delta^{13}\text{C}_{\text{CH}_4} > -49\text{‰}$ , approximately) are consistent with more than 60% removal of the deep endogenous component by bacterial oxidation, i.e., 40% of remaining deep endogenous  $\text{CH}_4$ . In cave air with the most depleted  $\text{CH}_4$  (<0.9 ppm and  $\delta^{13}\text{C}_{\text{CH}_4} > -35\text{‰}$ ), the percentage of remaining deep endogenous  $\text{CH}_4$  ranges 20–30%. In the case of the outstanding concentrations of deep endogenous  $\text{CH}_4$ , registered during March of 2016, it is demonstrated that the in situ  $\text{CH}_4$  oxidation process was not strong enough to deplete the upwelling flux of this gas below the atmospheric background and, consequently, the percentage of the remaining deep endogenous  $\text{CH}_4$  range is above 70%.

The presence of scattered data pairs diverging from the modelled distillation curves (consumption by MOB) in Figure 7 is due to that the closed-system Rayleigh model is an oversimplification of the  $\text{CH}_4$  dynamic. It assumes that the upwelling flux of deep endogenous methane is

not influenced by other potential inputs and the observed isotopic composition is not affected by other postgenetic processes as isotopic fractionation by diffusion or molecular fractionation by advection. However, oxidation of  $\text{CH}_4$  by MOB and the postgenetic modifications seem to occur simultaneously as the upwelling flux of endogenous air travels along the cave profile.

Isotopic fractionation by gas diffusion (curve [1] in Figure 7) is generated during the slow gas movement driven by concentration gradients. The result is a depletion of  $^{13}\text{C}$  in diffusing  $\text{CH}_4$  (corresponding data pairs diverging from the modelled distillation curves) and  $^{13}\text{C}$  enrichment in the residual gas (preferably corresponding to the rest of cave air samples with heavier  $\delta^{13}\text{C}_{\text{CH}_4}$ ). The isotopic fractionation by diffusion generally leads to a slight difference in  $\delta^{13}\text{C}_{\text{CH}_4}$ , not exceeding 5‰ [53]. This agrees with the  $\delta^{13}\text{C}_{\text{CH}_4}$  measurements for the air samples assigned as a result of this process (ranging from 1 to 30 m deep), which does not decrease below  $-56\text{‰}$  (Figure 7), i.e., 4.12‰, less than  $\delta^{13}\text{C}_{\text{CH}_4}$  for the maximum  $\text{CH}_4$  concentration registered at 50 m deep in March 2016 and considered as a clear example of deep endogenous source. In principle, a mixing process between cave air (with depleted  $\text{CH}_4$  due to methanotrophic activity) and a potential biogenic source (with

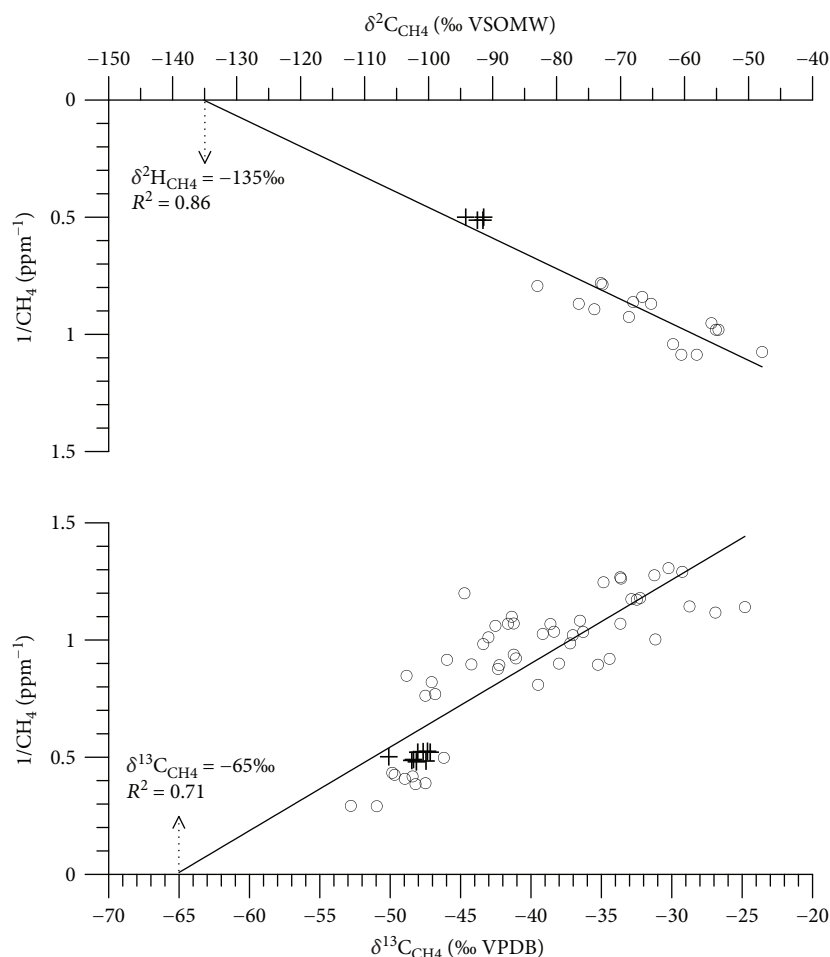


FIGURE 8: Keeling plots of  $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$  from cave air. The open circles represent cave air samples, and crosses represent the outdoor atmosphere. The Keeling plot of  $\delta^{13}\text{C}_{\text{CH}_4}$  was made with the entire set of air samples collected throughout the eight field campaigns, whereas the  $\delta^2\text{H}_{\text{CH}_4}$ -Keeling was plotted with the measurements obtained during the last two field campaigns. The vertical dotted arrows indicate the isotopic compositions ( $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$ ) for the source of  $\text{CH}_4$  that has been added to the cave air to produce the composition observed at each point.

$\delta^{13}\text{C}_{\text{CH}_4}$  roughly  $-65\text{‰}$ , according to Figure 8) is discarded because these data points of residual  $\text{CH}_4$  with lighter  $\delta^{13}\text{C}_{\text{CH}_4}$  do not fit properly with the hypothetical mixing curve between both end-members.

Molecular fractionation by advection is a sort of distillation likely provoked during the vertical transport of the endogenous warm air and upwelling along the cave profile. This process could be responsible for the differential segregation of light  $\text{CH}_4$  (data pairs with  $\delta^{13}\text{C}_{\text{CH}_4} < 60\text{‰}$  that fit curve [2] suggested in Figure 7), and it is exclusively observed for the air samples collected at 2 m depth, i.e., exhaled air from the cave to the open atmosphere.

An alternative mechanism for the isotopic fractionation by gas diffusion may be the low-temperature ( $<100^\circ\text{C}$ ) synthesis of  $\text{CH}_4$  related to gas-water-rock reactions, occurring in geothermal areas in continental settings and, even, at shallow depths. Several experimental studies have shown that abiotic  $\text{CH}_4$  derived by gas-water-rock reactions can result in  $\delta^{13}\text{C}_{\text{CH}_4}$  values as depleted as  $-57\text{‰}$  ([54, 55], and references therein), comparable to the isotopically light values

observed in VC and assigned to the isotopic fractionation by gas diffusion (curve [1] at Figure 7). In this sense, a potential inorganic mechanism for  $\text{CH}_4$  generation is the hydrogenation of  $\text{CO}_2$  in the gas phase (range of temperatures:  $25\text{--}500^\circ\text{C}$ , according to [54]) and the  $\text{H}_2$  necessary for this reaction could be produced by radiolytic decomposition of water vapour ( $\text{H}_2\text{O}_v$ ) due to the intense radioactive decay. Some of these conditions meet in the subterranean atmosphere of VC: geothermal activity ( $>35^\circ\text{C}$ ), large  $\text{CO}_2$  contents ( $>1\%$ ), and high environmental radioactivity ( $^{222}\text{Rn}$  higher than  $50\text{ kBq/m}^3$ ). Therefore, further research based on monitoring other ancillary gases as  $\text{H}_2$  is essential to providing better insights concerning the potential production of  $\text{CH}_4$  related to gas-water-rock reactions that do not directly involve organic matter.

For the next step of data analysis aimed at recognizing the geochemical features of the  $\text{CH}_4$  source, the set of  $[\text{CH}_4, \delta^{13}\text{C}_{\text{CH}_4}]$  data pairs identified as a likely result of the aforementioned postgenetic physical processes occurring in the cave profile, i.e., data pairs diverging from the modelled

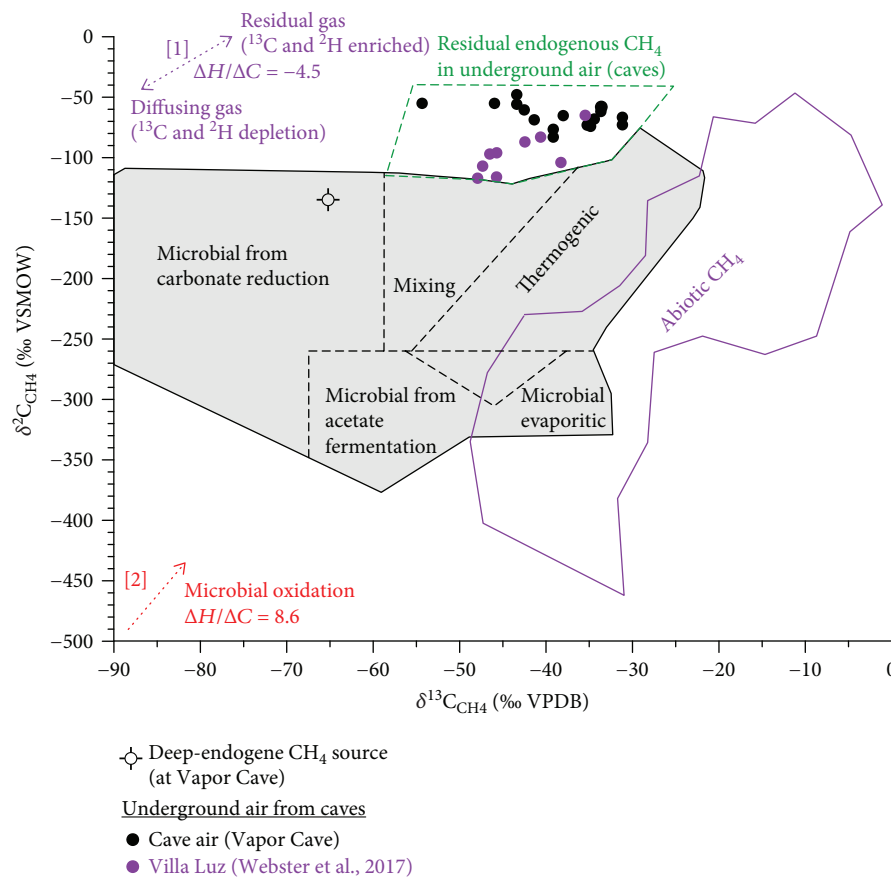


FIGURE 9: Genetic zonation of  $\text{CH}_4$  based on the isotopic composition of carbon ( $^{13}\text{C}/^{12}\text{C}$ ) and hydrogen ( $^2\text{H}/^1\text{H}$ ). Proposed genetic zones of biotic  $\text{CH}_4$  (grey zones): thermogenic, microbial from carbonate reduction, microbial from acetate fermentation, and microbial evaporitic, compared to the genetic zone for abiotic  $\text{CH}_4$  (modified from [54]). Data pairs of  $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$  are plotted for the local atmosphere, soil air, and underground air of VC in comparison with other published data from hypogenic acid caves [7]. Arrows correspond to two potential postgenetic processes affecting the molecular and isotopic composition of  $\text{CH}_4$  after its formation and during its migrations: [A] microbial oxidation and [B] gas diffusion.

distillation curves due to  $\text{CH}_4$  consumption by MOB, was not considered (Figure 7).

The relationships between  $\text{CH}_4$  concentration and  $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$  values, in both cave air and local atmosphere, fit a two end-member mixing model in Keeling plots (Figure 8).  $\delta^{13}\text{C}_{\text{CH}_4}$  in cave air ranged from  $-53\text{‰}$  to  $-25\text{‰}$ , with a subset of points quite similar to the local atmospheric  $\text{CH}_4$ .  $\delta^2\text{H}_{\text{CH}_4}$  ranged from  $-83\text{‰}$  to  $-48\text{‰}$ , i.e., with values markedly heavier than the local atmospheric background. The isotopic composition of source  $\text{CH}_4$  ( $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$ ) is estimated with both Keeling plots by extrapolating the linear function that fits the set of data points of cave air and local atmosphere, as far as its intersection with the  $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$  axis, respectively. Accordingly, the isotopic composition estimated for the source that contributes to  $\text{CH}_4$  content in cave air is roughly  $-65\text{‰}$  ( $\delta^{13}\text{C}_{\text{CH}_4}$ ) and  $-135\text{‰}$  ( $\delta^2\text{H}_{\text{CH}_4}$ ); i.e., this would correspond to the pure  $\text{CH}_4$  that is originally added to the upwelling air that reaches the cave environment and then it is consumed by MOB, independently of other postgenetic alteration processes affecting the concentration and isotopic composition of  $\text{CH}_4$  after its formation and emanation from this source

(e.g., isotopic fractionation by gas diffusion or molecular fractionation by advection).

The comparative analysis of the stable carbon and hydrogen isotope compositions of methane is an essential diagnostic tool to infer the origin of this gas, even though some additional interpretative parameters are needed for a better understanding, e.g., isotopic composition of associated gases as  $\text{CO}_2$ . The genetic zonation of  $\text{CH}_4$  based on the isotopic composition of carbon ( $^{13}\text{C}/^{12}\text{C}$ ) and hydrogen ( $^2\text{H}/^1\text{H}$ ) was originally introduced by [56] and then developed by [47, 54], among other authors. This analysis approach is aimed at distinguishing the specific signature of biotic methane (thermogenic and microbial) from other potential and diverse abiotic origins of gas (mainly due to volcanic/geothermal activity), besides to infer any sign of postgenetic alteration processes occurring before air is collected into the cave.

Figure 9 shows a diagram of the genetic zonation of  $\text{CH}_4$  with a well-defined distribution of carbon and hydrogen isotopes, based on worldwide occurrences of biotic and abiotic methane studied and revised by [54]. The isotopic range of  $\text{CH}_4$  observed in VC has been plotted in this diagram,



including local atmosphere, soil air, and underground air. The isotopic composition of the  $\text{CH}_4$  source inferred by Keeling plot analyses indicates that methane is primarily formed by bacterial carbonate reduction, likely linked to the Triassic black dolostones below the cave and under an intense water-rock interaction, according to the local hydrogeology settings described above. This kind of biotic methane is typically depleted in  $^{13}\text{C}$  relative to thermogenic and other biotic processes and usually range from  $<-100\text{‰}$  to about  $-50\text{‰}$  [47].

The stable isotopic composition of  $\text{CH}_4$  analysed for cave air does not closely match known isotopic compositions typical of microbial (biotic) or abiotic generation and is driven largely by relatively heavier  $\delta^2\text{H}_{\text{CH}_4}$  values. As a reference, methane in air of VC is even more  $^2\text{H}$ -enriched than  $\text{CH}_4$  recently described in active acid-hypogenic caves, e.g., Villa Luz [7] (Figure 9). This fact indicates that upwelling  $\text{CH}_4$  that reaches VC is the remnant of a larger  $\text{CH}_4$  flux at a depth that has been altered during its migration. Therefore, the gas sampled in the cave environment is clearly different from the original gas at the source, whose isotopic features have been previously inferred by Keeling analysis ( $\delta^{13}\text{C}_{\text{CH}_4}$ :  $-65\text{‰}$  and  $\delta^2\text{H}_{\text{CH}_4}$ :  $-135\text{‰}$ ). The partial consumption of  $\text{CH}_4$  and the associated shift in its isotopic compositions contents are likely due to the combination of two postgenetic secondary processes: a prevailing microbial oxidation and, in a less extent, an isotopic fractionation by diffusion. Both processes have been already brought up and inferred by analysing the relationship between  $\text{CH}_4$  and  $\delta^{13}\text{C}_{\text{CH}_4}$  in underground air, but now they are also corroborated with the comparative analysis of the  $\delta^{13}\text{C}_{\text{CH}_4}$  and  $\delta^2\text{H}_{\text{CH}_4}$  values. Microbial oxidation imparts an increase of about 8.5% in the  $\delta^2\text{H}_{\text{CH}_4}$  values for every increase of 1% in  $\delta^{13}\text{C}_{\text{CH}_4}$  [57, 58]. Gas diffusion, driven by concentration gradients according to Fick's law, may produce  $^{13}\text{C}$  and  $^2\text{H}$  depletion in the diffusing gas, and a residual gas will be  $^{13}\text{C}$ - and  $^2\text{H}$ -enriched, according to a  $\Delta H/\Delta C$  fractionation slope of 4.5 [54].

The isotopic composition of the deep endogenous source of  $\text{CH}_4$  and both fractionation slopes (microbial oxidation and gas diffusion) can be used to infer the alteration pathway of the upwelling  $\text{CH}_4$  at VC. Thus, the postgenetic alteration of biotic  $\text{CH}_4$  is a consequence of a simultaneous or sequenced effect of both microbial oxidation and gas diffusion. Both processes entail the production of residual methane that is  $^{13}\text{C}$ - and  $^2\text{H}$ -enriched, and it is primarily present in cave air, whereas the lighter  $\text{CH}_4$  (not usually sampled) is either consumed by MOB in a high percentage or is part of diffusing gas exhaled by the cave to the open atmosphere.

## 7. Conclusions

The gas composition of the subterranean atmosphere at VC is dominantly controlled by the upwelling airflow from the zone of fluid-geodynamic influence of active faulting. Data mining and modelling of variations in the concentrations of the main deep endogenous gases ( $\text{CO}_2$ ,  $\text{CH}_4$ ) and their isotopic signatures ( $\delta^{13}\text{C}_{\text{CO}_2}$  and  $\delta^{13}\text{C}_{\text{CH}_4}$ ) have provided considerable insight into the nature of gas exchange between

the atmospheric, soil, and underground air gas reservoirs at VC.

Degassing from  $\text{CO}_2$ -rich groundwater and deep-sourced geothermal  $\text{CO}_2$  (mantle-rooted gas) determine the high abundance of this gas at VC ( $>1\%$ ), with a heavier carbon isotopic composition, ranging from  $-4.5$  to  $-7.5\text{‰}$ .  $\text{CO}_2$  in underground air is typically composed of more than 30% of the pure theoretical  $\text{CO}_2$  added from a deep endogenous source. The cave acts as a net emitter of  $\text{CO}_2$  gas to the local atmosphere, so the exhaled air represents between 1 and 3% of this pure theoretical  $\text{CO}_2$  added from a deep endogenous source. The upwelling flow of deep endogenous air also provokes an intense  $\text{CO}_2$  diffusion from the cave air into the upper soil layers through fissures and small-size cracks in deep soil-epikarst. Thus, the diffusing  $\text{CO}_2$  measured for some soil air samples represent between 5% and 10% of the original deep endogenous  $\text{CO}_2$  sourcing the cave environment.

The source of methane in VC has an isotopic signature, which was likely generated by microbial carbonate reduction, likely affecting the Triassic black dolostones below the cave where it is the groundwater level of the local aquifer and, consequently, the water-rock interaction is higher. In this study, we have provided the first evidence demonstrating that caves may occasionally act as net sources of deep endogenous  $\text{CH}_4$  to the open atmosphere, with concentrations above the atmospheric background (ranging 2.3–3.4 ppm). This biotic  $\text{CH}_4$  is progressively oxidized during its migration through the upper vadose zone. Finally, subatmospheric concentrations of  $\text{CH}_4$  were registered in the cave environment. Therefore, hypogenic cave environments may also play a key role in regulating the release of greenhouse gases (e.g.,  $\text{CH}_4$ ) to the lower troposphere, through depletion of the concentration of methane with a deep endogenous origin.

## Data Availability

The raw data of  $\text{CO}_2$  and  $\text{CH}_4$  concentrations and their stable isotopic compositions used to support the findings of this study are available from the corresponding author upon request.

## Conflicts of Interest

There are no conflicts of interest to declare.

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